Box Sea 07/646

PATENT

Attorney Docket No.: A-68294/DJB/RMS/DAV

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of:

LUO et al.

Serial No. 09/404,010

Filed: September 23, 1999

For: NOVEL TRAF4 ASSOCIATED

CELL CYCLE PROTEINS,

COMPOSITIONS AND METHODS

OCT 1 0 2000

OF USE

Examiner: Andres, Janet

Group Art Unit: 1646

OCT 13 COM

CERTIFICATE OF MAILING

Thereby certify that this correspondence, including listed enclosures, is being deposited with the United States Postal Service as First Class Mail in an envelope addressed to: Assistant Commissioner for Patents, Washington, DC 2021 or:

Dated:

Cianad

TRANSMITTAL OF RESPONSE TO COMMUNICATION

AND NOTICE TO COMPLY

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Assistant Commissioner for Patents Washington, DC 20231

OCT 16 2000

TECH CENTER 1600/2900

Sir:

In response to the Office Communication and accompanying Notice to Comply with Requirements for Patent Applications Containing Nucleotide Sequence and/or Amino Acid Sequence Disclosures dated September 5, 2000, enclosed are the following documents:

Copy of Communication and Notice to Comply;

Serial No.: 09/404,010 **Filed**: September 23, 1999



- 2. Replacement Paper copy of SEQUENCE LISTING;
- 3. Communication Re: Sequence Listing
- 4. Replacement Computer readable copy of Sequence Listing.

While no fee is currently believed to be due, the Commissioner is authorized to charge any additional fees, including any extension fees, which may be required, or credit any overpayment to Deposit Account No. 06-1300 (Our Order No. A-68294/DJB/RMS/DAV).

Respectfully submitted,

FLEHR HOHBACH TEST ALBRITTON & HERBERT LLP

Dated: 10-4-00

Dolly A. Vance Reg. 39,054

Four Embarcadero Center Suite 3400 San Francisco, CA 94111-4187 Telephone: (415) 781-1989

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UNITED STATEDEPARTMENT OF COMMERCE Patent and Trademark Office

Address: COMMISSIONER OF PATENTS AND TRADEMARKS Washington, D.C. 20231

APPLICATION NO. FILING DATE FIRST NAMED INVENTOR ATTORNEY DOCKET NO.

09/404,010

09/23/99

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A-68294/DJB/

HM22/0905

FLEHR HOHBACH TEST ALBRITTON & TEST LLP FOUR EMBARCADERO CENTER STE 3400

SAN FRANCISCO CA 94111-4187

OLI 1 0 2000 4

EXAMINER

ANDRES.J

ART UNIT

PAPER NUMBER

1646

DATE MAILED:

09/05/00

Please find below and/or attached an Office communication concerning this application or proceeding.

' Commissioner of Patents and Trademarks

File A-68294 Atty RM5/DAV
Due Date 10|5|00 3/5/01

Type Zisting Refs

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OCT 16 2000



UNITED STATES DEPARTMENT OF COMMERCE Patent and Trademark Office COMMISSIONER OF PATENTS AND TRADEMARKS Washington, D.C. 20231

SERIAL NUMBER	FILING DATE	FIRST NAMED APPLICANT	ATTORNEY DOCKET NO.
09/404010	9/23/99	Luo	A68294/DJB/

EXAMINER				
Janet Andres				
ART UNIT	PAPER NUMBER			
1646	8			
DATE MAILED:				

Please find below a communication from the EXAMINER in charge of this application

Commissioner of Patents

- 1. This application contains sequence disclosures that are encompassed by the definitions for nucleotide and/or amino acid sequences set forth in 37 CAR 1.821(a)(1) and (a)(2). However, this application fails to comply with the requirements of 37 CAR 1.821 through 1.825 for the reason(s) set forth on the attached Notice To Comply With Requirements For Patent Applications Containing Nucleotide Sequence And/Or Amino Acid Sequence Disclosures.
- 2. Applicant is given ONE MONTH, or THIRTY DAYS, whichever is longer, from the mailing date of this letter within which to comply with the sequence rules, 37 CAR 1.821 1.825. Failure to comply with these requirements will result in ABANDONMENT of the application under 37 CAR 1.821(g). Extensions of time may be obtained by filing a petition accompanied by the extension fee under the provisions of 37 CAR 1.136(a). In no case may an applicant extend the period for reply beyond the SIX MONTH statutory period. Direct the reply to the undersigned. Applicant is requested to return a copy of the attached Notice to Comply with the reply.
- 3. Any inquiry concerning this communication or earlier communications from the examiner should be directed to Janet Andres, Ph.D. whose telephone number is (703)305-0557. The examiner can normally be reached on Monday-Thursday from 8:00-5:30 pm (Eastern Time) and 8:00-4:30 PM every other Friday.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Yvonne Eyler, Ph.D., can be reached at (703)308-6564. The FAX phone number for group 1600 is (703)308-4242.

An inquiry of a general nature or relating to the status of the application should be directed to the group receptionist whose telephone number is (703)308-0196.

YVONNE EYLER, PH.D PRIMARY EXAMINER

Application	.: <u>.</u>	09	<i> </i> 40	<u>401</u>	

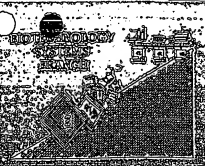
NOTICE TO COMPLY WITH REQUIREMENTS FOR PATENT APPLICATIONS CONTAINING NUCLEOTIDE SEQUENCE AND/OR AMINO ACID SEQUENCE DISCLOSURES

Applicant must file the items indicated below within the time period set the Office action to which the Notice is attached to avoid abandonment under 35 U.S.C. § 133 (extensions of time may be obtained under the provisions of 37 CFR 1.136(a)).

The nucleotide and/or amino acid sequence disclosure contained in this application does not comply with the requirements for such a disclosure as set forth in 37 C.F.R. 1.821 - 1.825 for the following reason(s):

X	1. This application clearly fails to comply with the requirements of 37 C.F.R. 1.821-1.825. Applicant's attention is directed to the final rulemaking notice published at 55 FR 18230 (May 1, 1990), and 1114 OG 29 (May 15, 1990). If the effective filing date is on or after July 1, 1998, see the final rulemaking notice published at 63 FR 29620 (June 1, 1998) and 1211 OG 82 (June 23, 1998).
	2. This application does not contain, as a separate part of the disclosure on paper copy, a "Sequence Listing" as required by 37 C.F.R. 1.821(c).
	3. A copy of the "Sequence Listing" in computer readable form has not been submitted as required by 37 C.F.R. 1.821(e).
X	4. A copy of the "Sequence Listing" in computer readable form has been submitted. However, the content of the computer readable form does not comply with the requirements of 37 C.F.R. 1.822 and/or 1.823, as indicated on the attached copy of the marked -up "Raw Sequence Listing."
	5. The computer readable form that has been filed with this application has been found to be damaged and/or unreadable as indicated on the attached CRF Diskette Problem Report. A Substitute computer readable form must be submitted as required by 37 C.F.R. 1.825(d).
	6. The paper copy of the "Sequence Listing" is not the same as the computer readable from of the "Sequence Listing" as required by 37 C.F.R. 1:821(e).
	7. Other: Sequences in application that do not have a sequence id number
Аp	pplicant Must Provide:
X	An initial or substitute computer readable form (CRF) copy of the "Sequence Listing".
X	An initial or <u>substitute</u> paper copy of the "Sequence Listing", as well as an amendment directing its entrinto the specification.
X	A statement that the content of the paper and computer readable copies are the same and, where applicable, include no new matter, as required by 37 C.F.R. 1.821(e) or 1.821(f) or 1.821(g) or 1.825(b) or 1.825(d).
Fo	or questions regarding compliance to these requirements, please contact:
Fo	or Rules Interpretation, call (703) 308-4216 or CRF Submission Help, call (703) 308-4212 otentIn Software Program Support
	Technical Assistance703-287-0200 To Purchase PatentIn Software703-306-2600
	PLEASE RETURN A COPY OF THIS NOTICE WITH YOUR REPLY

OHEN CHILISHING



The Biotechnology Systems Branchiof the Scientific and Technical Information Genter (STIC) detected errors when processing the following computer readable

Application Serial Number 29/40/50/0

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Source

Date Processed by STIC

TECH CENTER 1600/2

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.

PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY eitheer

- DENCLUDING A COPY OF THIS PRINTOUT IN YOUR WEXT COMMUNICATION TO THE APPLICANT WITH AMORIGE TO COMPLY
- 2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT WITH A NOTICE TO COMPLY
- FOR FURTHER INFORMATION PLEASE TELEPHONE MARK SPENCER 703-308-2212

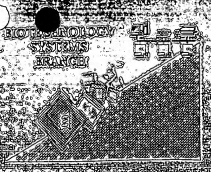
TO REDUCE ERRORED SEQUENCE LISTINGS REEASE USE THE CHECKE <u>VERSION 3:0 PROGRAM</u> ACCESSIBLE THROUGH THE UIS PATENT AND TRADEMARK OFFICE WEBSITE SEE BELOW

The Checker Version 3.0 application is a state of the art Windows based software program employing a logical and intuitive user interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listing generated for the original version of 37/CFR 661821 = 1825 effective October 1/1990 (old rules) and the revised version (new rules) effective ruly 181998 as well as a world intellectual Property Organization (WIPO) Standard ST:2

Checker Version 3:0 replaces the previous DOS-based version of Checker and is Y2K compliant Checker allows public users to check sequence listings in Computer Readable from F (CRF) before submitting them to the United States Patent and II rademark Office (USETO) is a Use of Checker prior coaling the sequence listing as expected to result in temer emoral sequence istines, thus saving time and money!

Thecker Wersion & O. san be down loaded from the USTAD website an the following address: harday and hardio-constrebiolaices (paralchealer

<u>ALW SIEGITEN ĈE LISTUM</u> C PERROR REPORT



The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) detected errors when processing the following computer readable

Application Serial Number: 09/405070

Source: 16.46

Date Processed by STIC: 7-26-00

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TECH CENTER 1600/200

THE ATTACHED PRINTOUT EXPLAINS DETECTED ERRORS.

PLEASE FORWARD THIS INFORMATION TO THE APPLICANT BY EITHER

- 1) INCLUDING A COPY OF THIS PRINTOUT IN YOUR NEXT COMMUNICATION TO THE APPLICANT, WITH A NOTICE TO COMPLY
- 2) TELEPHONING APPLICANT AND FAXING A COPY OF THIS PRINTOUT WITH A NOTICE TO COMPLY

FOR FURTHER INFORMATION, PLEASE TELEPHONE MARK SPENCER 703-308-4212

TO REDUCE ERRORED SEQUENCE LISTINGS PLEASE USE THE CHECKER VERSION 3:0 PROGRAM, ACCESSIBLE THROUGH THE U.S PATENT AND TRADEMARK OFFICE WEBSITE SEE BELOW

Checker Version 3.0

The Checker Version 3.0 application is a state-of the-art Windows based software program employing a logical and intuitive user-interface to check whether a sequence listing is in compliance with format and content rules. Checker Version 3.0 works for sequence listings. generated for the original version of 37. CFR §§1.821 = 1.825 effective October 1. 1990 (old rules) and the revised version (new rules) effective July 1: 1998 as well as World Intellectual Property Organization (WIPO) Standard ST-25

Checker Version 3:0 replaces the previous DOS-based version of Checker, and is V2Kcompliant. Checker allows public users to check sequence listings in Computer-Readable form is (CRF) before submitting them to the United States Patent and Trademark Office (USPTO) Use of Checker prior of filing the sequence listing is expected to result in fewer errored sequence listings thus saving time and money.

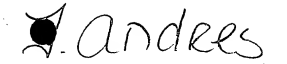
heckeraversion Biolizan bedown loaded from the USPIO website at the tollowing address x

The we we was a confer of the on the

hittor/Arvytuuspio-gov/Areb/oiffices/pac/checke

ERROR DETECTED SUGGESTED CORRECTION SERIAL NUMBER: 09/409

ATTN		LEASE DISREGARD ENGLISH "ALPHA" HEADERS, WHICH WERE INSERTED BY PTO SOFTWARE	
1	Wrapped Nucleics	The number/text at the end of each line "wrapped" down to the next line. This may occur if your file was retrieved in a word processor after creating it. Please adjust your right margin to .3, as this will prevent "wrapping".	VFL
2	Wrapped Aminos	The amino acid number/text at the end of each line "wrapped " down to the next line. This may occur if your file was retrieved in a word processor after creating it. AUG 16	200u
_		TECH CENTER 18	300/290 0
3	Incorrect Line Length	The rules require that a line not exceed 72 characters in length. This includes spaces.	
4	Misaligned Amino Acid Numbering	The numbering under each 5th amino acid is misaligned. This may be caused by the use of tabs between the numbering. It is recommended to delete any tabs and use spacing between the numbers.	<u>:</u> .
5	Non-ASCII	This file was not saved in ASCII (DOS) text, as required by the Sequence Rules. Please ensure your subsequent submission is saved in ASCII text so that it can be processed.	
6	Variable Length	Sequence(s) contain n's or Xaa's which represented more than one residue. As per the rules, each n or Xaa can only represent a single residue. Please present the maximum number of each residue having variable length and Indicate in the (ix) feature section that some may be missing.	
7	Patentin ver. 2.0 "bug"	A "bug" in PatentIn version 2.0 has caused the <220>-<223> section to be missing from amino acid sequence(s) Normally, PatentIn would automatically generate this section from the previously coded nucleic acid sequence. Please manually copy the relevant <220>-<223> section to the subsequent amino acid sequence. This applies primarily to the mandatory <220>-<223> sections for Artificial or Unknown sequences.	
8	Skipped Sequences (OLD RULES)	Sequence(s) missing. If intentional, please use the following format for each skipped sequence: (2) INFORMATION FOR SEQ ID NO:X: (i) SEQUENCE CHARACTERISTICS:(Do not insert any headings under "SEQUENCE CHARACTERISTIC (xi) SEQUENCE DESCRIPTION:SEQ ID NO:X: This sequence is intentionally skipped	(CS")
		Please also adjust the "(iii) NUMBER OF SEQUENCES:" response to include the skipped sequence(s).	
9	Skipped Sequences (NEW RULES)	Sequence(s) missing. If Intentional, please use the following format for each skipped sequence. <210> sequence id number <400> sequence id number 000	
10 💉	Use of n's or Xaa's (NEW RULES)	Use of n's and/or Xaa's have been detected in the Sequence Listing. Use of <220> to <223> is MANDATORY if n's or Xaa's are present. In <220> to <223> section, please explain location of n or Xaa, and which residue n or Xaa represents.	
l1	Use of <213>Organism (NEW RULES)	Sequence(s) are missing this mandatory field or its response.	
I2	Use of <220>Feature (NEW RULES)	Sequence(s) are missing the <220>Feature and associated headings. Use of <220> to <223> is MANDATORY if <213>ORGANISM is "Artificial" or "Unknown" Please explain source of genetic material in <220> to <223> section. (See "Federal Register," 6/01/98, Vol. 63, No. 104, pp. 29631-32) (Sec. 1.823 of new	w Rules)
13	Patentin ver. 2.0 "bug"	Please do not use "Copy to Disk" function of Patentin version 2.0. This causes a corrupted file, resulting in missing mandatory numeric identifiers and responses (as indicated on raw sequence listing instead, please use "File Manager" or any other means to copy file to floopy disk)). ·



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RAW SEQUENCE LISTING

Input Set : A:\A682941.app

PATENT APPLICATION: US/09/404,010

Output Set: N:\CRF3\07262000\I404010.raw

DATE: 07/26/2000

TIME: 14:25:18

Does Not Comply Corrected Diskette Needed

3 <110> APPLICANT: Luo, Ying Xu, Xiang 71 <210> SEQUENCES

71 <210> SEQ ID NO: 2
72 <211> LENGTH: 832
73 <212> TYPE RNA
74 <213> ORGANISM: Homo sapiens
76 <400> SEQUENCE: 2
77 nsgnnaEEaP gakaber
78 atgspur
78 atgspur
78 atgspur
78 atgspur
78 6 <120> TITLE OF INVENTION: Novel Traf4 Associated Cell Cycle Proteins, ERRORED SEQUENCES sequence contains E--> 77 nsgnnaEEaP gakaPEPaaa vothwFFard PvrdFPFELI PEPPEggLPg PwaLhrgrkk 60 E--> 78 atgsPvsIFv ydvkPgaEEQ @vakaaFkr FktLrhPnIL ayldgLEtEk cLhvvtEavt 120 "n's". As per E--> 79 PLgIyLkarv EaggLkELEI swgLhQIvka LsFLvndcsL Ihnnvcmaav FvdragEwkL 180 E--> 80 ggLdymysaQ gngggPPrkg IPELEQydPP ELadssgrvv rEkwsadmwr LgcLIwEvFn 240 E--> 81 gPLPraaalr nPgkIPktLa PhycELvgan PkvrPnParF LQncraPggF msnrFvEthL 300 E--> 82 FLEEIQIKEP aEkQkFFQEL sksLdaFPEd FcrhkLLPQL LtaFEFgnag avvLtPLFkv 360 sequence, rules, E--> 83 gkFLsaEEyQ QkIIPvvvkm FsstdramrI rLLQQmEQFI QyLdEPtvnt QIFPhvvhgF 420 E--> 84 IdtnPaIrEQ tvksmLLLaP kLnEanLnvE LmkhFarLQa kdEQgPIr(n)ttvcLgkIgs 480 E--> 85 yLsastrhrv Ltsafsratr dPFaPsrvag vLgFaathnL ysmndcaQkI LPvLcgLtvd 540 E--> 86 PEksvrdQaF kaFrsFLskL EsvsEdPtQL EEvEkdvhaa ssPgmggaaa swagwavtgv 600 "n's" most be E--> 87 ssLtskLIrs hPttaPtEtn IPQrPtPEgv PaPaPtPvPa tPttsghwEt QEEdkdtaEd 660 E--> 88 sstadrwddE dwgsLEQEaE svLaQQddws tggQvsrasQ vsnsdhkssk sPEsdwsswE 720 E--> 89 aEgswEQgwQ EPssQEPPsd gtrLasEynw ggPEssdkgd PFatLsarPs tQPrPdswgE 780 E--> 90 dnwEgLEtds rQvkaELark krEErrrEmE akraErkvak gPmkLgarkL dZ addressed in Letters in upper case are (220) to (223) not valid bose identifiers. feature. * See #10 09

Error Summary Sheet VERIFICATION SUMMARY

PATENT APPLICATION: US/09/404,010

DATE: 07/26/2000 TIME: 14:25:19

Input Set : A:\A682941.app

Output Set: N:\CRF3\07262000\1404010.raw

```
L:77 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:77 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
L:77 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:77 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
L:77 M:340 W: (46) "n" or "Xaa" used: Feature required, for SEQ ID#:2
L:77 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:25
L:77 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:1
L:77 M:112 C: (48) String data converted to lower case,
L:78 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:78 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
L:78 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:78 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
M:340 Repeated in SeqNo=2
L:78 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:26
L:78 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:7
M:112 Repeated in SeqNo=2
L:79 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
 L:79 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
L:79 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
 L:79 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
 L:79 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:21
 L:79 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:7
 L:80 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
 L:80 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
 L:80 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:80 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
 L:80 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:20
 L:80 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:7
L:81 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
 L:81 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
 L:81 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:81 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
 L:81 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:24
L:81 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:9
L:82 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:82 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
 L:82 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:82 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
 L:82 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:36
L:82 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:11
 L:83 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
 L:83 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
 L:83 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:83 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
  L:83 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:31
  L:83 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:14
 L:84 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:84 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
```

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OCT 16 2000

VERIFICATION SUMMARY

10

DATE: 07/26/2000 TIME: 14:25:19

PATENT APPLICATION: US/09/404,010

Input Set : A:\A682941.app

Output Set: N:\CRF3\07262000\I404010.raw

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L:84 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:84 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
L:84 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:27
L:84 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:18
L:85 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:85 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
L:85 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:85 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:85 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
L:85 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:20
L:85 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:23
L:86 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:19
L:86 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:25
L:87 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:87 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
L:87 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:87 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
 L:87 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:35
 L:87 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:37
 L:88 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
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L:89 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
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L:89 M:258 W: Mandatory Feature missing, <222> not found for SEQ ID#:2
L:89 M:258 W: Mandatory Feature missing, <223> not found for SEQ ID#:2
L:89 M:300 F: (1) Wrong Musleic Acid Designator, NUMBER OF INVALID PROCESS.
  L:89 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:25
L:89 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:42
  L:90 M:258 W: Mandatory Feature missing, <220> not found for SEQ ID#:2
L:90 M:258 W: Mandatory Feature missing, <221> not found for SEQ ID#:2
  L:90 M:320 E: (1) Wrong Nucleic Acid Designator, NUMBER OF INVALID KEYS:16
  L:90 M:321 E: (1) "t" not allowed in RNA Sequence, NUMBER OF INVALID 't' KEYS:43
```

AUG 10

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